A 2 OIPE

RAW SEQUENCE LISTING DATE: 09/25/2001 PATENT APPLICATION: US/09/847,809 TIME: 14:43:02

Input Set : N:\Crf3\RULE60\09847809.txt
Output Set: N:\CRF3\09252001\1847809.raw

## SEQUENCE LISTING

```
C--> 5 (1) GENERAL INFORMATION:
             (i) APPLICANT: Bandman, Olga
      7
                            Hillman, Jennifer L.
      8
      9
                            Lal, Preeti
     10
                            Corley, Neil C.
                            Shah, Purvi
     11
            (ii) TITLE OF INVENTION: HUMAN RETICULOCALBIN ISOFORMS
C--> 13
           (iii) NUMBER OF SEQUENCES: 6
     15
            (iv) CORRESPONDENCE ADDRESS:
     17
                  (A) ADDRESSEE: Incyte Pharmaceuticals, Inc.
     18
     19
                  (B) STREET: 3174 Porter Drive
     20
                  (C) CITY: Palo Alto
     21
                  (D) STATE: CA
                  (E) COUNTRY: USA
     22
     23
                  (F) ZIP: 94304
                                                            ENTERED
             (v) COMPUTER READABLE FORM:
     25
                  (A) MEDIUM TYPE: Diskette
     26
     27
                  (B) COMPUTER: IBM Compatible
                  (C) OPERATING SYSTEM: DOS
     28
                  (D) SOFTWARE: FastSEQ for Windows Version 2.0
     29
     31
            (vi) CURRENT APPLICATION DATA:
C--> 32
                  (A) APPLICATION NUMBER: US/09/847,809
                  (B) FILING DATE: 01-May-2001
C--> 33
                  (C) CLASSIFICATION:
     34
           (vii) PRIOR APPLICATION DATA:
     36
     37
                  (A) APPLICATION NUMBER: 09/270,270
                  (B) FILING DATE: 1999-03-16
     38
          (viii) ATTORNEY/AGENT INFORMATION:
     40
     41
                  (A) NAME: Billings, Lucy J.
                  (B) REGISTRATION NUMBER: 36,749
     42
                  (C) REFERENCE/DOCKET NUMBER: PF-0358 US
     43
            (ix) TELECOMMUNICATION INFORMATION:
     45
                  (A) TELEPHONE: 650-855-0555
     46
                  (B) TELEFAX: 650-845-4166
     47
     48
                  (C) TELEX:
        (2) INFORMATION FOR SEQ ID NO: 1:
     51
             (i) SEQUENCE CHARACTERISTICS:
     53
     54
                  (A) LENGTH: 328 amino acids
     55
                  (B) TYPE: amino acid
                  (C) STRANDEDNESS: single
     56
                  (D) TOPOLOGY: linear
     57
     59
           (vii) IMMEDIATE SOURCE:
                  (A) LIBRARY: RATRNOT02
     60
                  (B) CLONE: 922578
     61
     63
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
        Met Met Trp Arg Pro Ser Val Leu Leu Leu Leu Leu Leu Leu Arg His
```

Input Set : N:\Crf3\RULE60\09847809.txt
Output Set: N:\CRF3\09252001\I847809.raw

66	1				5					10					15	
67		Δla	Gln	Glv	_	Pro	Ser	Pro	Δsn		Glv	Pro	His	Glv		Gly
68	Gry	nια	OIII	20	цуз	110	JCI	110	25	1114	017	110		30	01	011
69	Δra	Va 1	Hic		Δla	Δla	Pro	T.e.11		Asn	Δla	Pro	His		Asp	Ala
70	лгу	Val	35	OIII	AIG	niu	110	40	DCI	пор		110	45	1156	шър	1114
71	Hic	Clv		Dhe	Gln	туг	Asn		Glu	Δla	Phe	T.e.u		Ara	Glu	Val
72	1113	50	ADII	1110	0.111	- 7 -	55	1115	Olu	11.1.4	1	60	011	9	014	, 42
73	717		Clu	Dha	λen	Gln		Thr	Dro	Glu	Glu		Gln	Δla	Arg	T.e.11
74	65	пуз	GIU	FIIC	пор	70	пси	1111	110	GIU	75	001	0111	mu	**** 9	80
75		7 ~~	Tla	Wa l	Nen		Mot	λen	) ra	λla		Aen	Cl v	Aen	Gly	
76	Gry	AIG	110	Val	85	nrg	ricc	пор	n. 9	90	OLY	nsp	013	mpp	95	
77	Val	Ser	T.e.u	Δla		Len	Δrσ	Δla	Ψrn		Δla	His	Thr	Gln	Gln	Ara
78	Val	SCI	шец	100	Olu	БСи	ni 9	niu	105	110	ri_u	1115	****	110	01	
79	Wi c	Tlo	λνα		Car	V ⊃ l	Cor	בוג		Ψrn	Aen	Thr	ጥላን		Thr	Asn
80	птр	116	115	изр	361	Val	361	120	Ата	пр	пэр	1111	125	пор	1111	Nop
81	7 ~~	7 an		λνα	Wa l	Clv	Птр		Glu	T.011	λνα	Acn		Thr	Tyr	Glv
	AIG	_	GLY	AIG	vaı	GLY	135	GIU	GIU	пец	лгу	140	AIG	1111	- <u>y</u> -	GLY
82	TT-1 -	130	71-	Dwo	C1	~1.v		Dha	uio	7 cr	Val.		7 cn	7 l n	Glu	Thr.
83		туг	Ald	PIO	GTÄ		GIU	Pne	птъ	ASP	155	GIU	ASP	Ата	Giu	160
84	145	T	T	Wat	T 0.11	150	7 ~~	7 00	C1.,	7 ~~		Dho	7 ~~	17 n 1	λla	
85	TYL	гаг	гаг	Met		Ala	AIG	ASP	GIU	170	Arg	Pile	AIG	Val	Ala 175	кър
86	G1	3	a1	7 ~~	165	Wot	7.1.	Пhm	7 ~~		C1.,	T 011	Mhr	717		T All
87	GIN	ASP	GIĀ	_	ser	Met	Ala	THI	_	GIU	GIU	Leu	TIIT	190	Phe	цеu
88	***	D		180	Dh.	D	114.	Mat	185	7 0 0	т1 о	17 n 1	Tlo		C1.,	mh.r.
89	HIS	Pro	195	GIU	Pne	Pro	HIS		Arg	Asp	TTE	val	205	Ald	Glu	TIII
90	*	<b>~1</b>		т	7 an	7 ~~	200	200	N an	C1.,	Птт	W-1		17 n 1	Clu	Clu
91	Leu		Asp	ьeu	ASP	Arg	215	цуѕ	ASP	GLY	тут	220	GIII	Val	Glu	Giu
92	m	210	71-	7 00	T 011	Птт		7.1 -	Clu	Dro	C117		Clu	Glu	Pro	λla
93		TTE	Ald	ASP	ьeu	230	ser	нта	GIU	PIO	235	Giu	Giu	GIU	FIO	240
94 95	225	77a 1	~1 ~	Пhъ	C1		C1 n	Cln	Dho	7 ~~		Dho	7 ~~	7 cn	Leu	
96	тър	Val	GIII	TIII	245	ALG	GIII	GIII	FIIC	250	ASP	FIIC	Arg	кэр	255	ASII
96 97	T	7.00	C1	uio		7 an	C1,17	Cor	Clu		C1 17	uic	Πrn	Val	Leu	Pro
98	гуѕ	ASP	GIY	260	ьeu	АБР	GIY	ser	265	vaı	Gry	1112	ттр	270	шец	FIO
	Dwa	71.	C1 n		Cln	Dro	T 011	W-1		7.1.	7 an	ui.	Tou		Wic	Clu
99	Pro	Ald	27	_	GIII	PIO	пеп	280		нта	ASII	птъ	28		His	Giu
100	Co	~ 7 ~ 1			- T 17	- Acı	^ Cl;			, 60.	r T 177	- או			ь Т. <b>ь</b> і	u Gly
101	se.			LAS	5 гл.	s AS	29		у пе	ı se.	г гу.	30		u II.	e ne	u Giy
102	3	29		. Mad	- Dh	· 170			~ Cl,	. זו	- mъ			~ Cl	., Cl	u Asp
103			p ASI	i Me	L FIII			y 3e.	L GII	I AL			.1 ту.	L G1.	y Gi	
	104 305 310 315 320															
	105 Leu Thr Arg His His Asp Glu Leu															
	106 325 108 (2) INFORMATION FOR SEQ ID NO: 2:															
	(2)															
	110 (i) SEQUENCE CHARACTERISTICS:															
	111 (A) LENGTH: 1463 base pairs 112 (B) TYPE: nucleic acid															
112			-								1					
113							ESS:		эте							
114		,	•				line	=a I								
116		(V11	) IM					мошо.	2							
117			( 4	н) ш.	TDKA	KI;	RATRI	NOTU.	۷							

Input Set : N:\Crf3\RULE60\09847809.txt
Output Set: N:\CRF3\09252001\1847809.raw

```
(B) CLONE: 922578
118
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
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    CGCAGAGCGG ACGTGGAGAG CGGACGNCAG CTGGATAACA GGGGACCGAT GATGTGGCGA
                                                                           60
122
123 CCATCAGTTC TGCTGCTTCT GTTGCTACTG AGGCACGGGG CCCAGGGGAA GCCATCCCCA
                                                                          120
124 GACGCAGGCC CTCATGGCCA GGGGAGGGTG CACCAGGCGG CCCCCCTGAG CGACGCTCCC
                                                                          180
125 CATGATGACG CCCACGGGAA CTTCCAGTAC GACCATGAGG CTTTCCTGGG ACGGGAAGTG
                                                                          240
126 GCCAAGGAAT TCGACCAACT CACCCCAGAG GAAAGCCAGG CCCGTCTGGG GCGGATCGTG
    GACCGCATGG ACCGCGGGG GGACGGCGAC GGCTGGGTGT CGCTGGCCGA GCTTCGCGCG
                                                                          360
127
    TGGATCGCGC ACACGCAGCA GCGGCACATA CGGGACTCGG TGAGCGCGGC CTGGGACACG
128
                                                                          420
    TACGACACGG ACCGCGACGG GCGTGTGGGT TGGGAGGAGC TGCGCAACGC CACCTATGGC
                                                                          480
129
130 CACTACGCGC CCGGTGAAGA ATTTCATGAC GTGGAGGATG CAGAGACCTA CAAAAAGATG
                                                                          540
                                                                          600
131 CTGGCTCGGG ACGAGCGGCG TTTCCGGGTG GCCGACCAGG ATGGGGACTC GATGGCCACT
                                                                          660
132 CGAGAGGAGC TGACAGCCTT CCTGCACCCC GAGGAGTTCC CTCACATGCG GGACATCGTG
133 ATTGCTGAAA CCCTGGAGGA CCTGGACAGA AACAAAGATG GCTATGTCCA GGTGGAGGAG
134 TACATCGCGG ATCTGTACTC AGCCGAGCCT GGGGAGGAGG AGCCGGCGTG GGTGCAGACG
                                                                          780
135 GAGAGGCAGC AGTTCCGGGA CTTCCGGGAT CTGAACAAGG ATGGGCACCT GGATGGGAGT
                                                                          840
136 GAGGTGGGCC ACTGGGTGCT GCCCCCTGCC CAGGACCAGC CCCTGGTGGA AGCCAACCAC
                                                                          900
137 CTGCTGCACG AGAGCGACAC GGACAAGGAT GGGCGGCTGA GCAAAGCGGA AATCCTGGGT
                                                                          960
138 AATTGGAACA TGTTTGTGGG CAGTCAGGCC ACCAACTATG GCGAGGACCT GACCCGGCAC
                                                                         1020
139 CACGATGAGC TGTGAGCACC GCGCACCTGC CACAGCCTCA GAGGCCCGCA CAATGACCGG
140 AGGAGGGCC GCTGTGGTCT GGCCCCCTCC CTGTCCAGGC CCCGCAGGAG GCAGATGCAG
141 TCCCAGGCAT CCTCCTGCCC CTGGGCTCTC AGGGACCCCC TGGGTCGGCT TCTGTCCCTG
                                                                         1200
    TCACACCCC AACCCCAGGG AGGGGCTGTC ATAGTCCCAG AGGATAAGCA ATACCTATTT
142
                                                                         1260
143 CTGACTGAGT CTCCCAGCCC AGACCCAGGG ACCCTTGGCC CCAAGCTCAG CTCTAAGAAC
                                                                         1320
144 CGCCCCAACC CCTCCAGCTC CAAATCTGAG CCTCCACCAC ATAGACTGAA ACTCCCCTGG
                                                                         1380
145 CCCCAGCCCT CTCCTGCCTG GCCTGGCCTG GGACACCTCC TCTCTGCCAG GAGGCAATAA
                                                                         1440
146 AAGCCAGCGC CGGGAAAAAA AAA
                                                                         1463
148 (2) INFORMATION FOR SEQ ID NO: 3:
150
        (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 315 amino acids
151
152
              (B) TYPE: amino acid
153
              (C) STRANDEDNESS: single
              (D) TOPOLOGY: linear
    (vii) IMMEDIATE SOURCE:
156
157
              (A) LIBRARY: BLADNOT03
158
              (B) CLONE: 1601793 .
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
160
    Met Asp Leu Arg Gln Phe Leu Met Cys Leu Ser Leu Cys Thr Ala Phe
162
                      5
                                         10
163
    Ala Leu Ser Lys Pro Thr Glu Lys Lys Asp Arg Val His His Glu Pro
164
165
                 20
                                     25
     Gln Leu Ser Asp Lys Val His Asn Asp Ala Gln Ser Phe Asp Tyr Asp
166
167
                                 40
                                                     45
    His Asp Ala Phe Leu Gly Ala Glu Glu Ala Lys Thr Phe Asp Gln Leu
168
169
                                                 60
                             55
     Thr Pro Glu Glu Ser Lys Glu Arg Leu Gly Lys Ile Val Ser Lys Ile
170
                         70
                                             75
171
    Asp Gly Asp Lys Asp Gly Phe Val Thr Val Asp Glu Leu Lys Asp Trp
172
173
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Input Set: N:\Crf3\RULE60\09847809.txt
Output Set: N:\CRF3\09252001\1847809.raw

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Ile Lys Phe Ala Gln Lys Arg Trp Ile Tyr Glu Asp Val Glu Arg Gln
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175
                 100
                                     105
    Trp Lys Gly His Asp Leu Asn Glu Asp Gly Leu Val Ser Trp Glu Glu
176
                                 120
                                                     125
177
            115
178
     Tyr Lys Asn Ala Thr Tyr Gly Tyr Val Leu Asp Asp Pro Asp Pro Asp
                             135
179
    Asp Gly Phe Asn Tyr Lys Gln Met Met Val Arg Asp Glu Arg Arg Phe
180
                                             155
                         150
181
    Lys Met Ala Asp Lys Asp Gly Asp Leu Ile Ala Thr Lys Glu Glu Phe
182
                                         170
183
                     165
    Thr Ala Phe Leu His Pro Glu Glu Tyr Asp Tyr Met Lys Asp Ile Val
185
186
                                     185
                 180
    Val Gln Glu Thr Met Glu Asp Ile Asp Lys Asn Ala Asp Gly Phe Ile
187
                                 200
188
     Asp Leu Glu Glu Tyr Ile Gly Asp Met Tyr Ser His Asp Gly Asn Thr
189
190
                             215
                                                  220
    Asp Glu Pro Glu Trp Val Lys Thr Glu Arg Glu Gln Phe Val Glu Phe
191
                         230
                                             235
192
    Arg Asp Lys Asn Arg Asp Gly Lys Met Asp Lys Glu Glu Thr Lys Asp
                                         250
194
                     245
     Trp Ile Leu Pro Ser Asp Tyr Asp His Ala Glu Ala Glu Ala Arg His
195
196
                 260
                                     265
197
    Leu Val Tyr Glu Ser Asp Gln Asn Lys Asp Gly Lys Leu Thr Lys Glu
198
             275
                                 280
    Glu Ile Val Asp Lys Tyr Asp Leu Phe Val Gly Ser Gln Ala Thr Asp
199
200
                             295
201
    Phe Gly Glu Ala Leu Val Arg His Asp Glu Phe
                         310
202
204 (2) INFORMATION FOR SEQ ID NO: 4:
        (i) SEQUENCE CHARACTERISTICS:
206
             (A) LENGTH: 2658 base pairs
207
208
            (B) TYPE: nucleic acid
209
              (C) STRANDEDNESS: single
              (D) TOPOLOGY: linear
210
       (vii) IMMEDIATE SOURCE:
212
              (A) LIBRARY: BLADNOT03
213
              (B) CLONE: 1601793
214
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
216
218 CCCGCTTCCG GTTGGGCGGT GCTTGCGCGC GTGAGCTGAG CCGGTGGGTG AGCGGCGGCC
                                                                            60
                                                                           120
    ACGGCATCCT GTGCTGTGGG GGCTACGAGG AAAGATCTAA TTATCATGGA CCTGCGACAG
    TTTCTTATGT GCCTGTCCCT GTGCACAGCC TTTGCCTTGA GCAAACCCAC AGAAAAGAAG
                                                                           180
220
    GACCGTGTAC ATCATGAGCC TCAGCTCAGT GACAAGGTTC ACAATGATGC TCAGAGTTTT
221
     GATTATGACC ATGATGCCTT CTTGGGTGCT GAAGAAGCAA AGACCTTTGA TCAGCTGACA
                                                                           300
222
    CCAGAAGAG GCAAGGAAAG GCTTGGAAAG ATTGTAAGTA AAATAGATGC CGACAAGGAC
                                                                           360
223
    GGGTTTGTCA CTGTGGATGA GCTCAAAGAC TGGATTAAAT TTGCACAAAA GCGCTGGATT
                                                                           420
    TACGAGGATG TAGAGCGACA GTGGAAGGGG CATGACCTCA ATGAGGACGG CCTCGTTTCC
                                                                           480
    TGGGAGGAGT ATAAAAATGC CACCTACGGC TACGTTTTAG ATGATCCAGA TCCTGATGAT
                                                                           540
    GGATTTAACT ATAAACAGAT GATGGTTAGA GATGAGCGGA GGTTTAAAAT GGCAGACAAG
227
                                                                           600
    GATGGAGACC TCATTGCCAC CAAGGAGGAG TTCACAGCTT TCCTGCACCC TGAGGAGTAT
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Input Set : N:\Crf3\RULE60\09847809.txt
Output Set: N:\CRF3\09252001\1847809.raw

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229 GACTACATGA AAGATATAGT AGTACAGGAA ACAATGGAAG ATATAGATAA GAATGCTGAT
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230 GGTTTCATTG ATCTAGAAGA GTATATTGGT GACATGTACA GCCATGATGG GAATACTGAT
                                                                          780
    GAGCCAGAAT GGGTAAAGAC AGAGCGAGAG CAGTTTGTTG AGTTTCGGGA TAAGAACCGT
231
                                                                          840
232
    GATGGGAAGA TGGACAAGGA AGAGACCAAA GACTGGATCC TTCCCTCAGA CTATGATCAT
                                                                          900
233 GCAGAGGCAG AAGCCAGGCA CCTGGTCTAT GAATCAGACC AAAACAAGGA TGGCAAGCTT
234 ACCAAGGAGG AGATCGTTGA CAAGTATGAC TTATTTGTTG GCAGCCAGGC CACAGATTTT
                                                                         1020
235 GGGGAGGCCT TAGTACGGCA TGATGAGTTC TGAGCTACGG AGGAACCCTC ATTTCCTCAA
236 AAGTAATTTA TTTTTACAGC TTCTGGTTTC ACATGAAATT GTTTGCGCTA CTGAGACTGT
                                                                         1140
    TACTACAAAC TTTTTAAGAC ATGAAAAGGC GTAATGAAAA CCATCCCGTC CCCATTCCTC
237
                                                                         1200
    CTCCTCTCT AGGGACTGGA GGGAAGCCGT GCTTCTGAGG AACAACTCTA ATTAGTACAC
238
                                                                         1260
    TTGTGTTTGT AGATTTACAC TTTGTATTAT GTATTAACAT GGCGTGTTTA TTTTTGTATT
239
                                                                         1320
240 TTTCTCTGGT TGGGAGTATG ATATGAAGGA TCAAGATCCT CAACTCACAC ATGTAGACAA
                                                                         1380
241 ACATTAGCTC TTTACTCTTT CTCAACCCCT TTTATGATTT TAATAATTCT CACTTAACTA
                                                                         1440
242 ATTTTGTAAG CCTGAGATCA ATAAGAAATG TTCAGGAGAG AGGAAAGAAA AAAAATATAT
243 GCTCCACAAT TTATATTTAG AGAGAGAACA CTTAGTCTTG CCTGTCAAAA AGTCCAACAT
                                                                         1560
    TTCATAGGTA GTAGGGGCCA CATATTACAT TCAGTTGCTA TAGGTCCAGC AACTGAACCT
                                                                         1620
244
    GCCATTACCT GGGCAAGGAA AGATCCCTTT GCTCTAGGAA AGCTTGGCCC AAATTGATTT
                                                                         1680
246 TCTTCTTTTT CCCCCTGTAG GACTGACTGT TGGCTAATTT TGTCAAGCAC AGCTGTGGTG
                                                                         1740
247
    GGAAGAGTTA GGGCCAGTGT CTTGAAAATC AATCAAGTAG TGAATGTGAT CTCTTTGCAG
                                                                         1800
248 AGCTATAGAT AGAAACAGCT GGAAAACTAA AGGAAAAATA CAAATGTTTT CGGGGCATAC
    ATTTTTTTC TGGGTGTGCA TCTGTTGAAA TGCTCAAGAC TTAATTATTT GCCTTTTGAA
249
    ATCACTGTAA ATGCCCCCAT CCGGTTCCTC TTCTTCCCAG GTGTGCCAAG GAATTAATCT
250
                                                                         1980
    TGGTTTCACT ACAATTAAAA TTCACTCCTT TCCAATCATG TCATTGAAAG TGCCTTTAAC
251
                                                                         2040
    GAAAGAAATG GTCACTGAAT GGGAATTCTC TTAAGAAACC CTGAGATTAA AAAAAGACTA
252
                                                                         2100
253 TTTGGATAAC TTATAGGAAA GCCTAGAACC TCCCAGTAGA GTGGGGATTT TTTTCTTCTT
                                                                         2160
254 CCCTTTCTCT TTTGGACAAT AGTTAAATTA GCAGTATTAG TTATGAGTTT GGTTGCAGTG
                                                                         2220
255 TTCTTATCTT GTGGGCTGAT TTCCAAAAAC CACATGCTGC TGAATTTACC AGGGATCCTC
                                                                         2280
256 ATACCTCACA ATGCAAACCA CTTACTACCA GGCCTTTTTC TGTGTCCACT GGAGAGCTTG
    AGCTCACACT CAAAGATCAG AGGACCTACA GAGAGGGCTC TTTGGTTTGA GGACCATGGC
257
                                                                         2400
258 TTACCTTTCC TGCCTTTGAC CCATCACACC CCATTTCCTC CTCTTTCCCT CTCCCCGCTG
                                                                         2460
    CCAAAAAAA AAAAAAAGGA AACGTTTATC ATGAATCAAC AGGGTTTCAG TCCTTATCAA
                                                                         2520
    AGAGAGATGT GGAAAGAGCT AAAGAAACCA CCCTTTGTTC CCAACTCCAC TTTACCCATA
                                                                         2580
    TTTTATGCAA CACAAACACT GTCCTTTTGG GTCCCTTTCT TACAGATGGG ACCTCTTGAG
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    GAAGGAATTA TCGTATTC
                                                                         2658
264 (2) INFORMATION FOR SEQ ID NO: 5:
266
         (i) SEQUENCE CHARACTERISTICS:
267
              (A) LENGTH: 331 amino acids
268
              (B) TYPE: amino acid
269
             (C) STRANDEDNESS: single
270
             (D) TOPOLOGY: linear
272
       (vii) IMMEDIATE SOURCE:
273
              (A) LIBRARY: GenBank
274
              (B) CLONE: 1262329
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:
276
278
    Met Ala Arg Gly Gly Arg Gly Arg Leu Gly Leu Ala Leu Gly Leu
279
                      5
                                        10
280
    Leu Leu Ala Leu Val Leu Ala Pro Arq Val Leu Arq Ala Lys Pro Thr
                                     25
282 Val Arg Lys Glu Arg Val Val Arg Pro Asp Ser Glu Leu Gly Glu Arg
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VERIFICATION SUMMARY

DATE: 09/25/2001

PATENT APPLICATION: US/09/847,809

TIME: 14:43:03

Input Set : N:\Crf3\RULE60\09847809.txt Output Set: N:\CRF3\09252001\I847809.raw

L:5 M:220 C: Keyword misspelled or invalid format, [(1) GENERAL INFORMATION:] L:13 M:220 C: Keyword misspelled or invalid format, [(ii) TITLE OF INVENTION:]
L:32 M:220 C: Keyword misspelled or invalid format, [(A) APPLICATION NUMBER:]
L:33 M:220 C: Keyword misspelled or invalid format, [(B) FILING DATE:]